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(54) Title: OLIGONUCLEOTIDE REPEAT ARRAYS

(57) Abstract

A solid support based hybridization assay is provided which allows for the systematic and reproducible analysis of repeat and tandem repeat oligonucleotide sequences of DNA and RNA by hybridization to a reverse dot blot array comprising strings of such repeats complementary to those found in particular nucleic acid targets (e.g., analyte PCR product). An addressable library (i.e., an indexed set) of complementary repeats is synthesized on a suitable support. Preferably, the support comprises a low fluorescent background support, thereby facilitating the use of non-radioisotopic modes of detection (such as fluorescence of chemiluminescence); particularly suitable in this regard is an aminated polypropylene support or similar material. Preferred arrays permit screening of DNA and RNA samples for complete sets of particular types of nucleotide repeat sequences (e.g., all nucleotide doublet or triplet repeats).

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OLIGONUCLEOTIDE REPEAT ARRAYS

Background of the Invention

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The present invention relates generally to the fields of biochemistry and medicine. In particular, the invention is directed to materials and methods useful in the diagnosis of genetic mutations of clinical relevance.

Short tandem repeats (STR) have been identified in a number of genes. It has been proposed that particular unstable triplet repeat oligonucleotides are correlated with a number of genetic diseases in humans, including Kennedy's disease [La Spada, A. et al., Nature, 352, 77-79 (1991)], fragile-X syndrome [Verkerk, A.J.M.H. et al., Cell 65, 905-914 (1991)], myotonic dystrophy [Fu, Y.H. et al, Science 255, 1256-1258 (1992)], Huntington disease [The Huntington's Disease Collaborative Research Group, Cell 72, 971-983 (1993)] and spinocerebellar ataxia type 1 [Orr, H.T. et al., Nature Genet. 4, 221-226 (1993)]. Similarly, doublet repeats have also been reported to be associated with particular disease states; for example, correlations have been proposed with cystic fibrosis [Chu, C.-S. et al., Nature Genetics 3, 151-156 (1993)] and colorectal cancer [Thibodeau, S.N. et al., Science 260, 816-819 (1993)]. Higher-order repeats, such as tetramers [see, e.g., Gen, M.W. et al., Genomics 17, 770-772 (1993)], have also been identified.

One gene which has been subject of intense scrutiny is the Huntington's disease gene. The trinucleotide hybridization approach was recently utilized to map out tandem repeats across a section of the gene. section, 51 triplet repeats spanning a 1.86 Mbp DNA identified by Southern transfer were segment restriction enzyme digests of a specific cosmid and oligonucleotide 32P-labelled probing with [Hummerich, et al., "Distribution of trinucleotide repeat 2 Mbp region containing the sequences across a

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Huntington's disease gene, " Human Molecular Genetics 3, 73 (1994)].

DNA polymorphisms which arise from allelic differences in the number of repeats have been identified by such terminology as short tandem repeats (STR), variable number of tandem repeats (VNTR), minisatellites (tandem repeats of a short sequence, originally defined as 9-60 bp) and microsatellites (originally defined as 1-[McBride, L.J. & O'Neill, M.D., American Laboratory, pp. 52-54 (November 1991)]; minisatellites 10 and microsatellites would be considered subclasses of the It is estimated that there are up to 500,000 VNTR. microsatellite repeats distributed throughout the human genome, at an average spacing of 7000 bp. Therefore, it is apparent that most genes will contain VNTR regions and 15 that these regions can be used as genetic markers. For example, VNTRs are currently being used as markers in concerned with the inheritance of mutations leading to various forms of cancer. Recently, it has been discovered that certain triplet repeat 20 expansions are associated with a predisposition towards certain diseases; а large expansion is typically associated with the onset of the disease. For example, the (CGG) triplet repeat region associated with Fragile X occurs at a frequency of 10-50 repeat units in the 25 normal population, while in those afflicted with the disease the expansion is between 200-2000 repeats.

As it becomes possible to determine whether a particular genotype comprises an unstable repeat and/or is associated with a particular disease state, there is a considerable incentive to develop useful methods to characterize STRs. The heretofore available methods for initial scanning for STRs have generally required time-consuming sequential oligonucleotide hybridizations to filter-bound target DNAs to identify specific STRs [see, e.g., Litt, M. and Luty, J.A., Am. J. Hum. Genet. 44, 397-401 (1989); Weber, J.L. and May, P.E., Am. J. Hum.

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Genet. 44, 388-396 (1989); Fu et al., supra]. In particular, the analysis of oligonucleotide repeats is typically carried out at the present time by Southern blotting of restriction fragments followed by hybridization analysis using a specified repetitive sequence probe. Alternatively, it is possible to probe dot blots of the target DNA [Iizuka, et al., GATA 10:2-5 (1993)].

Both of these heretofore-known techniques are time-consuming and tedious for large sample populations. Moreover, multiple probings may be required to identify which repeat might be present. Further, it is often difficult to reproducibly spot or transfer equivalent amounts of DNA to these supports; thus, conventional dot blots and transfers show variation in signal intensity from batch to batch. In addition, any regions of DNA that might become cross-linked to the support (e.g., through UV light) would be inaccessible to probes.

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It would be highly useful for clinical investigators to be able to screen large sample populations of patients DNAs in an effective manner. As additional STRs are identified and associated with particular conditions, the need for simple and effective screening methods becomes greater.

PCT published application No. WO 89/10977 describes 25 methods and apparatus for analyzing polynucleotide sequences in which an array of the whole or a chosen part of a complete set of oligonucleotides are bound to a solid support. The different oligonucleotides occupy separate cells of the array and are capable of taking 30 hybridization reactions. For studying in differences between polynucleotide sequences, the array may comprise the whole or a chosen part of a complete set oligonucleotides comprising the polynucleotide sequences. While it is suggested that a small array may 35 be useful for many applications, such as the analysis of a gene for mutations, there is no teaching or suggestion of a specific array or method for using same which would permit the rapid and accurate screening of a wide range of biological materials for tandem repeats. Moreover, the arrays described in WO 89/10977 are designed specifically for use in sequencing by hybridization; the presence of long tandem nucleotide repeats can present a significant problem in attempts to sequence a sample using the methods described in WO 89/10977.

It is an object of the present invention to provide 10 methods and apparatus for rapid and accurate identification of nucleotide tandem repeats in DNA and RNA sequences from a wide variety of sources.

Summary of the Invention

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In accordance with the present invention, a solid support based hybridization assay is provided which allows for the systematic and reproducible analysis of repeat and tandem repeat oligonucleotide sequences of DNA and RNA by hybridization to a reverse dot blot array comprising strings of such repeats complementary to those found in particular nucleic acid targets (e.g., analyte PCR product). An addressable library (i.e., an indexed set) of complementary repeats is synthesized on a Preferably, the support comprises a suitable support. low fluorescent background support, thereby facilitating the use of non-radioisotopic modes of detection (such as fluorescence or chemiluminescence); particularly suitable in this regard is an aminated polypropylene support or similar material. Pursuant to a preferred embodiment of the invention, arrays are provided which permit screening of DNA and RNA samples for complete sets of particular all sequences (e.g., repeat of nucleotide types nucleotide doublet or triplet repeats).

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Brief Description of the Drawings

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The invention may be better understood with reference to the accompanying drawings, in which:

Fig. 1 illustrates a pattern for synthesis of leader sequences and tandem repeat patterns;

Figs. 2(A) - 2(C) illustrate hybridization of arrayed dinucleotide and trinucleotide oligonucleotide repeats using ³²P labelled DNA of cosmid 22.3 as a probe, in which Fig. 2(A) represents hybridization stringency at 40°C, Fig. 2(B), hybridization stringency at 50°C, and Fig. 2(C), hybridization stringency at 60°C; and

Fig. 3 illustrates the type and position of STRs indicated by the STR-Strips in 34,977 bp of cosmid 22.3.

Detailed Description of the Invention

In accordance with the present invention, defined repeat and tandem repeat arrays for use in screening nucleic acid targets for the presence of genetic markers generally known as variable number [of] tandem repeats (VNTRs) are synthesized on a suitable support. After hybridization of the target materials with the array, the identity of any tandem repeat sequence(s) in the target materials may be readily ascertained by observing the location(s) at which binding occurs. Pursuant to the present invention, probes are reproducibly synthesized on the surface, freely accessible to target DNA. Moreover, all hybridizations can be rapidly identified under a limited number of stringency conditions.

The arrays of the present invention comprise a predetermined set of oligonucleotides attached to the surface of the solid support. One particularly useful class of tandem repeats for arrays in accordance with the present invention comprises the complete class of 60 tandem triplet repeats (i.e., all possible triplet combinations minus the four homopolymer combinations).

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Another useful class of tandem repeats is the complete class of 6 tandem doublet repeats (i.e., the 10 possible doublet combinations of the nucleic acids A, C, G and T minus the four homopolymer combinations). Of course, those skilled in the art would readily appreciate that a . wide range of different combinations of repeat elements could also be employed in accordance with the present invention. For example, repeats of a higher order (i.e., repeats of four or more nucleotides) may be useful in some instances. In addition, particular subclasses of 10 any complete class of all possible tandem repeats of a given size may be suitable for carrying out particular types of screenings For purposes of the present invention, all predetermined sets of tandem repeats are scope of the contemplated as within the 15 invention.

The sequences forming the array may be directly In other embodiments of the linked to the support. arrays of the present invention, the repeat units may be attached to the support by non-repetitive sequences of oligonucleotides or other molecules serving as spacers or linkers to the solid support. In preferred examples of this embodiment, specific leader sequences are encoded on either side of the tandem repeat region in an array Depending upon the relative position of the format. leader sequence a PCR or sequencing primer may be Such primers may then be used to aid in the characterization of the length of the tandem repeat and/or the specific flanking sequences, respectively. general, a triplet tandem repeat sequence of sufficient length effectively defines two additional tandem repeat sequences; for example, a 21mer complementary to (ACG)_n also hybridizes to (GAC), and (CGA),. By systematically including a degenerate set of leader sequences while the tandem repeat reducing the size of hybridization stringency is increased to allow for identification of the combination of leader plus the

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tandem repeat; in the example, selectivity of CCC ACG ACG ACG [SEQ ID NO:1] would be observed over, e.g., CCC GAC GAC GAC [SEQ ID NO:2]. Fig. 1 illustrates a set of instructions for synthesis of suitable leader sequences for triplet tandem repeats. Such leader arrays are particularly advantageous for the purpose of identifying leader sequences for use as PCR primers to tandem repeat regions.

The method of the present invention is generally applicable to a wide range of tandem repeat patterns, 10 including higher order tandem repeats. As by definition a tandem repeat consists of at least 2 units of a given oligomer (for example, a dimer or 2mer), then a (2mer), wherein n = 2, 3, 4, ... would represent a dinucleotide repeat forming a 4mer, 6mer, 8mer, etc. (e.g., ACAC, 15 Similarly, a triplet repeat ACACAC, ACACACAC, etc.). would be defined as a (3mer), and a tetramer repeat as (4mer), wherein n represents the number of repeats present. Contemplated as within the scope of the present invention are all tandem repeats of the general formula 20 integer greater wherein N is an representing the number of nucleotides in the repeat pattern and n is an integer representing the number of times the pattern is repeated; in general, the product of N and n is in the range of 4 to about 100, and preferably 25 6 to about 60.

Higher order tandem repeat combinations representing combinations of two or more individual tandem repeats are also contemplated as within the scope of the present invention; for example, such higher order tandem repeat combinations may include two dimer patterns, a dimer and a triplet, two triplets, etc. In general terms, such repeat combinations may be described as

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(Nmer), (Mmer),

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independently selected integers and represent the number of times the respective pattern is repeated. In the case of three tandem patterns, the structure may be represented as

(Nmer) (Mmer) (Pmer) p

in which P is defined in the same manner as N and M, and p in the same manner as n and m. Moreover, these higher order tandem repeat combinations may also be found in a repeat pattern; such a complex higher order tandem repeat combination may be described as

[(Nmer),(Mmer),)]x

or

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[(Nmer)_n(Mmer)_m(Pmer)_p]_x

in which N, M, P, n, m and p are as previously defined and x is an integer which represents the number of times the [(Nmer)_n(Mmer)_m)] or [(Nmer)_n(Mmer)_m(Pmer)_p] pattern is repeated. Contemplated as within the scope of the present invention are those combinations wherein x(Nn + Mm) or x(Nn + Mm + Pp) is in the range of 4 to about 100, preferably in the range of 6 to about 60. In the simplest case comprising two repeat patterns and wherein x is 1, N and M are both 2 and n and m are both 1; 1 x 2 + 1 x 2 = 4. Table 1 illustrates the construction of higher order repeats in which n and m are both 2.

25 TABLE 1

Tandem Repeat	(2mer)	(3mer)	(4mer) _n	(5mer) _n	(6mer) _n	(7mer) _n
(2mer) _m	8mer	10mer	12mer	14mer	16mer	18mer
(3mer) _m	10mer	12mer	14mer	16mer	18mer	20mer
(4mer) _m	12mer	14mer	16mer	18mer	20mer	22mer
(5mer) _m	14mer	16mer	18mer	20mer	22mer	24mer
(6mer) _m	16mer	18mer	20mer	22mer	24mer	26mer
(7mer) _m	18mer	20mer	22mer	24mer	26mer	28mer

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Thus, for example, the 8mer comprising a $(2mer)_n = (AC)_2$ and a $(2mer)_m = (AT)_2$ would have the formula ACACATAT;

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similarly, the 12mer comprising a $(3mer)_n = (ACG)_2$ and a $(3mer)_m = (ATC)_2$ would have the formula ACGACGATCATC [SEQ ID NO:3]. A complex higher order repeat $[(AC)_n(AT)_m]_x$ in which n and m are 1 and x is 2 would have the formula ACATACAT; where x is 3, the formula would be ACATACATACAT [SEQ ID NO:4]. Those skilled in the art would readily appreciate the variety of simple and higher-order tandem repeat patterns falling within the scope of the present invention.

10 The methods and apparatus in accordance with the present invention take advantage of the fact that under appropriate conditions oligonucleotides form base paired duplexes with oligonucleotides which have a complementary The stability of the duplex is dependent base sequence. on a number of factors, 15 including the length of the oligonucleotides, composition, the base composition of the solution in which hybridization is The effects of base composition on duplex effected. carrying stability may be reduced by hybridization in particular solutions, for example in the 20 presence of high concentrations of tertiary or quaternary amines.

The thermal stability of the duplex is also dependent on the degree of sequence similarity between the sequences. By carrying out the hybridization at temperatures close to the anticipated Tm's of the type of duplexes expected to be formed between the target material(s) and the oligonucleotides bound to the array, the rate of formation of mismatched duplexes may be substantially reduced.

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The number of repeats in the tandem sequences attached to the array may vary over a broad range from the minimum of two necessary to constitute a repeat to a maximum on the order of about 50. Of course, the optimum range for the number of tandem repeats in any given instance is dependent upon a number of factors, including in particular the composition and the length of the

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repeat. In general, the Tm of the complex formed between the target material sequence in complementary sequence in the array increases as the length of the sequences increase; however, only minor increases in Tm are observed once the sequences have reached a length of about 50-60 bases. The sequences on the array contain at least four bases (the minimum for a repeat of a doublet pattern). It is generally preferred that the sequences on the array comprise at least about 10 6 bases, more preferably at least about 10 bases, and most preferably on the order of about 15 to about 60 As previously noted, there is little practical advantage in using sequences substantially longer than about 60 bases; nonetheless, extended sequences of up to about 100 bases in length (corresponding to, e.g., 50 repeats of a doublet sequence) and longer are also scope of the present within the contemplated as invention.

with preferred in accordance In addition. embodiments of the invention the length of each sequence employed in the array may be selected to as to optimize binding of target materials to the array. For any given tandem repeat sequence, an optimum length for use with any particular target material under specified screening conditions may be determined empirically. Thus, the length for each individual element of the set of tandem repeats comprising the array may be optimized for the screening of particular target materials under specific conditions (e.g., at a given temperature).

A wide variety of array formats may be employed in accordance with the present invention. One particularly useful format is a linear array of oligonucleotide bands, generally referred to in the art as a dipstick. suitable format comprises a two-dimensional pattern of discrete cells (e.g., 4096 squares in a 64 by 64 array). as would be readily appreciated by those Of course, skilled in the art, other array formats (e.g., circular)

would be equally suitable for use in accordance with the While arrays may be prepared on a present invention. variety of materials including glass plates, presently preferred to use an organic polymer medium. used herein, the term "organic polymer" is intended to support material which is most preferably under conditions appropriate chemically inert biopolymer synthesis and which comprises a backbone comprising various elemental substituents including, but not limited to, hydrogen, carbon, oxygen, 10 chlorine, bromine, sulfur and nitrogen. Representative polymers include, but are not limited to, the following: polyethylene, polybutylene, polypropylene, polybutadiene, polyisoprene, polyisobutylene, polytetrafluoroethylene, polyvinylpyrrolidone, 15 polyvinylidene difluoride, polyfluoroethylene-propylene, alcohol, polymethylpentene, polyethylene-vinyl polychlorotrifluoroethylene, polysulfones, and blends and copolymers thereof. As used herein, the term "medium" is intended to mean the physical structural shape of the 20 Thus, medium can be generally defined as polymer films (i.e., polymers having a substantially nonporous surface); polymer membranes (i.e., polymers having a porous surface); polymer filaments (e.g., mesh and fabrics); polymer beads; polymer foams; polymer frits; 25 and polymer threads. Preferably, the polymer medium is a thread, membrane or film; most preferably, the polymer medium is a film. An exemplary organic polymer medium is a polypropylene sheet having a thickness on the order of about 1 mil (0.001 inch), although the thickness of the 30 film is not critical and may be varied over a fairly Particularly preferred for preparation of broad range. arrays at this time are biaxially oriented polypropylene (BOPP) films; in addition to their durability, BOPP films exhibit a low background fluorescence. 35

The array formats of the present invention may be included in a variety of different types of device. As

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used herein, the term "device" is intended to mean any device to which the solid support can be affixed, such as tubes, microtiter plates, test inorganic dipsticks, etc. For example, when the solid support is a polypropylene thread, one or more polypropylene threads can be affixed to a plastic dipstick-type device; polypropylene membranes can be affixed to glass slides. The particular device is, in and of itself, unimportant. All that is necessary is that the solid support can be affixed thereto without affecting the functional behavior of the solid support or any biopolymer adsorbed thereon, and that the device is stable to any materials into which the device is introduced (e.g., clinical samples, etc.).

The arrays of the present invention may be prepared by a variety of approaches which are known to those 15 working in the field. Pursuant to one type of approach, the complete sequences are synthesized separately and then attached to a solid support. However, it is presently considered particularly advantageous to synthesize the sequences directly onto the support to 20 Suitable methods provide the desired array. covalently coupling oligonucleotides to a solid support and for directly synthesizing the oligonucleotides onto the support would be readily apparent to those working in the field; a summary of suitable methods may be found in, 25 e.g., Matson, R.S. et al., Analytical Biochem. 217, 306reference. hereby incorporated by (1994), 310 Advantageously, the oligonucleotides are synthesized onto the support using conventional chemical techniques as heretofore employed for preparing oligonucleotides on 30 solid supports comprising, e.g., controlled pore size glass (CPG), as described for example in PCT applications WO 85/01051 and WO 89/10977, or polypropylene, described in copending U.S. patent application Serial No. 07/091,100, which has been assigned to the assignee of 35 the present application and is herein incorporated by Pursuant to one preferred approach, reference.

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polypropylene support (for example, a biaxially-oriented polypropylene) is first surface aminated by exposure to an ammonia plasma generated by radiofrequency plasma The reaction of a phosphoramidite-activated discharge. nucleotide with the aminated membrane followed oxidation with, e.g., iodine provides a base stable amidate bond to the support.

As described in U.S. patent application Serial No. 1993, filed October 28, which has been 08/144,954 commonly assigned to the assignee of the present 10 invention and is incorporated by reference herein, a suitable array may advantageously be produced using automated means to synthesize oligonucleotides in the cells of the array by laying down the precursors for the four bases in a predetermined pattern. Briefly, multiple-channel automated chemical delivery system is employed to create oligonucleotide probe populations in parallel rows (corresponding in number to the number of channels in the delivery system) across the substrate. Following completion of oligonucleotide synthesis in a 20 first (1°) direction, the substrate may then be rotated by 90° to permit synthesis to proceed within a second (2°) set of rows that are now perpendicular to the first set. This process creates a multiple-channel array whose intersection generates a plurality of discrete cells. 25 Table 2 describes an exemplary vertical array of 64 oligonucleotides consisting of 60 triplet tandem repeat sequences (21mers) and dinucleotide tandem repeat sequences (20mers).

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For the example of the preferred array comprising specific leader sequences as described in Fig. 1 (and as more fully described in the above-noted U.S. patent application Serial No. 08/144,954), all of the degenerate including homopolymers) (n=64,repeats synthesized in a first direction (1° synthesis) in the 64 35 channels in Cycles 1-3 as 3' Leader Sequences (LLL). example, lane 1 is AAA, lane 2 AAC, lane 5 ACA and lane 10

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64 TTT. Then, the film is rotated 90° to perform synthesis in the second direction (2° synthesis or crosssynthesis) in Cycles 4-15. All 64 triplet tandem repeat sequences of (NNN)₄ are then produced. The 2-dimensional array created thereby is the product of a bidirectional synthesis and comprises 4096 discrete cells containing 15mer oligonucleotide products (LLL)(NNN)₄, in which the Leader Sequence is placed at the 3'-end of each completed oligonucleotide. Thus, the following sequences would be found at the indicated cell positions:

Cell	1,1'	(AAA)	(AAA) ₄	[SEQ	ID	NO:5]
Cell	1,2'	(AAA)	(AAC) ₄	[SEQ	ID	NO:6]
Cell	5,1'	(ACA)	(AAA) ₄	[SEQ	ID	NO:7]
Cell	64,1'	(TTT)	(AAA) ₄	[SEQ	ID	NO:8]
Cell	64.64'	(TTT)	(TTT),	[SEO	ID	NO:9]

This type of array comprising leader sequences (at either the 5' or 3' end) is particularly preferred in accordance with the present invention.

In order to accommodate a suitably large array, the pixel size should be as small as possible. Cells having a width on the order of about 10 μm to about 1 mm would be particularly suitable. In one preferred embodiment of the invention, arrays with a pixel width of about 500 μm are prepared on biaxially-oriented polypropylene.

Pursuant to the present invention, there are also provided methods for screening DNA and RNA samples comprising labelling the samples to form material, applying the labelled material under suitable hybridization conditions to an array as described herein, and observing the location of the label on the surface associated with particular members of the oligonucleotides. Identification of the cell(s) in which rapid and accurate permits а binding occurs identification of any nucleotide repeats present in the sample from which the probes are derived.

In a hybridization reaction in accordance with the present invention, the array is explored by the labelled

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target material in essentially the same manner as a labelled probe is employed to screen, e.g., a DNA library containing a gene complementary to the probe. suitably comprise labelled sequences material may amplified from genomic DNA by the polymerase chain reaction (PCR), a mRNA population, or a partial or complete set of oligonucleotides from one chromosomes or an entire genome. To prepare the target material, the sample may be degraded to form fragments; where appropriate, the degraded material may then be sorted (for example, by electrophoresis on a sequencing gel) to provide a set of oligomers having a specific length.

The target material is then labelled to facilitate detection of duplex formation. Suitably, conventional 15 methods for end-labelling of oligomers are employed. Both radioactive and fluorescent labelling methods would be suitable for use in accordance with the present invention. Commonly-employed techniques routinely permit the introduction of label into a significant fraction of 20 Using conventional methods for the target materials. labelling oligomers with 32P, for example, the radioactive yield of any individual oligomer even from a total human genome could be more than 104 dpm/mg of total DNA. detection, only a small fraction of the labelled material would be necessary for hybridization to a pixel of a size specified preferred range within the Hybridization conditions for a given combination of array and target material can routinely be optimized in an empirical manner to be close to the Tm of the expected 30 duplexes, thereby maximizing the discriminating power of the method. Autoradiography (in particular, with 32P) may cause image degradation which may be a limiting factor determining resolution; the limit for silver halide films Accordingly, the use of fluorescent is about 25 μ m. 35 probes (in particular, in conjunction with an array low-fluorescence solid support) prepared on a

presently preferred. In view of the low background biaxially-oriented preferred fluorescence of the polypropylene substrate, fluorescence-based labelling techniques may advantageously be employed with arrays on With either type of labeled target such a substrate. substantial excess in bound material. the oligonucleotides of the array makes it possible to operate at conditions close to equilibrium with most types of target materials contemplated herein.

As would be readily understood by those skilled in 10 the art, the chosen conditions of hybridization must be such as to permit discrimination between exactly matched oligonucleotides. Hybridization mismatched conditions may be initially chosen to correspond to those suitable in standard procedures be 15 hybridization to filters and then optimized for use with the arrays of the present invention; moreover, conditions suitable for hybridization of one type of target material would appropriately be adjusted for use with other target materials for the same array. In particular, 20 appropriate to control temperature closely (preferably, to better than about ±1° C) to substantially eliminate formation of duplexes between sequences other than identical sequences. Particularly when the length of the oligonucleotide in the target materials is small, it is 25 necessary to be able to distinguish between slight differences in the rate and/or extent of hybridization.

A variety of heretofore known hybridization solvents may suitably be employed, the choice of solvent for particular hybridizations being dependent on a number of For example, G:C base pairs are more considerations. stable than A:T base pairs in 1 M NaCl; thus, the Tm of double-stranded oligonucleotides with a high G + C higher than corresponding will be content oligonucleotides with a high A + T content. particularly course pronounced in are of sequences comprising tandem nucleotide repeats.

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compensate for this discrepancy, a variety of to approaches may be employed. For example, the amount of oligonucleotide applied to the surface of the support may be varied in dependence on the nucleotide composition of the bound oligomer. Further, computer means employed to analyze data from hybridization experiments may for variations programmed to make compensations nucleotide compositions. Another expedient (which may be employed instead of or in addition to those already mentioned) is the use of a chaotropic hybridization . 10 solvent, such as a ternary or quaternary amine. tetramethylammonium chloride (TMAC1) at regard, concentrations in the range of about 2 M to about 5.5 M is particularly suitable; at TMACl concentrations around 3.5 to 4 M, the Tm dependence on nucleotide composition 15 In addition, the choice of is substantially reduced. hybridization salt has a major effect on overall hybridization yield; for example, TMACl at concentrations up to 5 M can increase the overall hybridization yield by a factor of up to 30 or more (depending to some extent on 20 nucleotide composition) compared to 1 M length of previously noted, the Finally, as oligonucleotides attached to the array may be varied so optimize hybridization under the particular conditions employed. As previously noted, it would be a 25 routine matter for those working in the field to optimize hybridization conditions for any given combination of target materials and array.

Hybridization is typically carried out with a very large excess of the bound oligonucleotides over what is 30 In preferred embodiments of the found in the target. invention, it is possible in some cases to distinguish between hybridization involving single and multiple sequence, as yield of the target occurrences proportional to concentration at all stages in the 35 reaction.

In accordance with another embodiment of the present invention, an array as described herein may be employed to selectively isolate and size variable number of tandem This is accomplished by preparing a repeats (VNTRs). sample comprising VNTRs in a manner known per se [see, e.g., McBride & O'Neill, supra], capturing the VNTRs on selectively dissociating the hybrid and the array, eluting the VNTR from the support. A selected lane from the array may be cut out of the support, the ssDNA eluted therefrom, the number of copies thereof increased by PCR amplification and size analysis conducted by conventional technique (e.g., gel electrophoresis against DNA size markers). The presence of large molecular weight strands would indicate an increase in mutational frequency (i.e., higher orders of tandem repeat regions).

The invention may be better understood with reference to the accompanying example that is intended for purposes of illustration only and should not be construed as in any sense limiting the scope of the present invention as defined in the claims appended hereto.

Example

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Oligonucleotides were synthesized directly from aminated 6.6 x 6.6 cm sheet of monomers onto а substrate using standard CEDpolypropylene A specially designed 64 phosphoramidite chemistries. channel chemical delivery system (Southern Array Maker™, Beckman Instruments) as described in co-pending U.S. patent application Serial No. 08/144,954 was utilized to oligonucleotide sequences prepare the discrete across the polypropylene substrate. parallel rows aminated surface by film was Polypropylene radiofrequency discharge into ammonia gas as described in The plasma-aminated film was then Matson et al., supra. placed in the synthesizer. Standard phosphoramidite chemistry was performed in each of the 64 channels to create 64 different oligonucleotide sequences on the

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film. The substrate was then cut into 0.5 cm widths perpendicular to the oligonucleotide rows to produce a panel of 64 tandem repeat sequences. For the present study 60 trinucleotide (21mers) and 4 dinucleotide tandem (20mers) repeat sequences were arrayed on a vertical order shown in Table 2. The arrayed trinucleotide repeat set represents all triplet frames except (AAA), [SEQ ID NO:10], (CCC), [SEQ ID NO:11], (GGG), [SEQ ID NO:12], and (TTT), [SEQ ID NO:13] in 3'-->5' direction as well as minus strand orientation.

In order to confirm that all sequences were fully represented on the panel a series of complementary probes were prepared that would verify each sequence by row position on the strip. As each triplet repeat (n) of a sufficient length in fact reads the n-1 and n-2 frames as well, only 20 probe sequences were required to identify the 60 triplet repeats on the panel. For example, row 2 containing the sequence (AAC), [SEQ ID NO:14] also represents (ACA) [SEQ ID NO:15] and (CAA) [SEQ ID NO:16] (sequences found at row 5 and row 17, respectively); thus, a single tandem repeat probe which has been labeled will identify the sequences at row positions 2, 5 and 17. Four additional probe sequences were required to identify the 4 doublet tandem repeats synthesized on the strip. It is also possible to combine the sequences of noncomplementary probes (i.e., those that will not selfhybridize or form hairpin loops) to reduce the total number of probes necessary to read all row positions. The following 32P 5'-end labeled probes were prepared and used to identify all row positions 1-64 (listed in Table 2) on the panel:

	(TGC) ₆ ;	[SEQ ID NO:17]
	$(TGG)_4(TG)_6(TTA)_4;$	[SEQ ID NO:18]
	(TGG) ₄ (TGA) ₄ ;	[SEQ ID NO:19]
35	(TCG) ₆ ;	[SEQ ID NO:20]
	(GGC) _s ;	[SEQ ID NO:21]
	(CG) ₄ (GAC) ₄ ;	[SEQ ID NO:22]

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$(TCA)_4(TAA)_4(TAC)_4;$	(SEQ	ID	NO:23]
(CCA) ₄ (CAA) ₄ ;	[SEQ	ID	NO:24]
(TTC) ₄ (TC) ₆ (TCC) ₄ ;	[SEQ	ID	NO:25]
(GAA) 4 (GA) 6 (GGA) 4;	and [SEQ	ID	NO:26]
(GCC),(GCA),	[SEO	ID	NO:27].

A number of different test DNAs were employed. This included a 5'-(CAG),-3' oligonucleotide [SEQ ID NO:28] and 200 bp PCR fragments containing a trinucleotide short tandem repeat of (CTG) 11 [SEQ ID NO:29] [Fu, Y.H., et al, Science 255, 1256-1258 (1992)] generated from human 10 An 800 bp PCR genomic DNA of a wild type individual. fragment containing a (CAG)₁₀ [SEQ ID NO:30] repeat and a 3.0 kb PCR fragment containing a tandemly combined (GCA)₈+(GCG)₄ repeat [SEQ ID NO:31] were generated from cDNA clones G13 and A12, respectively, recently isolated 15 in a new cDNA identification system [Lee, C.C., et al., Am. J. Hum. Genet. 53 (Suppl.), 1321 (1993)]. DNA samples of the cosmid MDY2 [Fu, Y.H., et al, Science 255, 1256-1258 (1992)] containing the entire myotonin protein kinase gene and cosmid 22.3 containing the FMR-1 gene 20 [Verkerk, A.J.M.H., et al., Cell 65, 905-914 (1991)] have also been utilized as target materials to evaluate the test strips.

STRs were identified using the GCG sequence analysis software package (Genetics Computer Group, Inc. 1991). 11,613 bp of cosmid MDY2 (GenBank accession L00727) and 61,612 bp of a contiguous sequence containing the complete 34,977 bp sequence of cosmid 22.3 were searched for STRs.

TABLE 2

Vertical array of 64 oligonucleotides consisting of 60 triplet tandem repeat sequences (21mers) and dinucleotide tandem repeat sequences (20mers) on a polypropylene substrate

5	#	Oligonucleotides 3' > 5'	Seq ID No.	#	Oligonucleotides 3' > 5'	Seq ID No.
	1	AC AC AC AC AC AC AC AC	32	33	GAA GAA GAA GAA GAA	62
	2	AAC AAC AAC AAC AAC	14	34	GAC GAC GAC GAC GAC	63
İ	3	AAG AAG AAG AAG AAG	33	35	GAG GAG GAG GAG GAG	64
10	4	TAA TAA TAA TAA TAA TAA	34	36	GAT GAT GAT GAT GAT	65
	5	ACA ACA ACA ACA ACA	35	37	GCA GCA GCA GCA GCA	66
	6	ACC ACC ACC ACC ACC ACC	36	38	GCC GCC GCC GCC GCC	67
į	7	ACG ACG ACG ACG ACG	37	39	GCG GCG GCG GCG GCG	68
	8	ACT ACT ACT ACT ACT ACT	38	40	GCT GCT GCT GCT GCT	69
15	9	AGA AGA AGA AGA AGA	39	41	GGA GGA GGA GGA GGA	70
ĺ	10	AGC AGC AGC AGC AGC	40	42	GGC GGC GGC GGC GGC	71
ŀ	11	AGG AGG AGG AGG AGG	41	43	AG AG AG AG AG AG AG AG	72
	12	AGT AGT AGT AGT AGT	42	44	GGT GGT GGT GGT GGT	73
	13	ATA ATA ATA ATA ATA ATA	43	45	GTA GTA GTA GTA GTA	74
20	14	ATC ATC ATC ATC ATC ATC	44	46	GTC GTC GTC GTC GTC	75
	15	ATG ATG ATG ATG ATG	45	47	GTG GTG GTG GTG GTG	76
	16	TTA TTA TTA TTA TTA TTA	46	48	GTT GTT GTT GTT GTT GTT	77
	17	CAA CAA CAA CAA CAA CAA	47	49	TAA TAA TAA TAA TAA TAA	78
	18	CAC CAC CAC CAC CAC CAC	48	50	TAC TAC TAC TAC TAC TAC	79
25	19	CAG CAG CAG CAG CAG CAG	28	51	TAG TAG TAG TAG TAG	80
	20	CAT CAT CAT CAT CAT CAT	49	52	TAT TAT TAT TAT TAT TAT	81
	21	CCA CCA CCA CCA CCA CCA	50	53	TCA TCA TCA TCA TCA TCA	82
i	22	CG CG CG CG CG CG CG CG	51	54	דכב דכב דכב דכב דכב דכב	83
	23	CCG CCG CCG CCG CCG	52	55	TCG TCG TCG TCG TCG TCG	84
30	24	CCT CCT CCT CCT CCT CCT	53	56	זכד זכד זכד זכד זכד זכד	85
	25	CGA CGA CGA CGA CGA	54	57	TGA TGA TGA TGA TGA	86
.	26	CGC CGC CGC CGC CGC	55	58	TGC TGC TGC TGC TGC	87
	27	CGG CGG CGG CGG CGG	56	59	TGG TGG TGG TGG TGG	88
	28	CGT CGT CGT CGT CGT	57	60	TGT TGT TGT TGT TGT TGT	89
35	29	CTA CTA CTA CTA CTA CTA	58	61	TTA TTA TTA TTA TTA TTA	90
	30	сте ете ете ете ете ете	59	62	דוכ זוכ זוכ דוכ דוכ דוכ דוכ	91
	31	CTG CTG CTG CTG CTG	60	63	דום דום דום דום דום דום דום	92
	32	כדד כדד כדד כדד כדד כדד	61	64	כד כד כד כד כד כד כד כד כד	93

Oligonucleotide probes were end-labelled with 32P-Gamma-dATP under standard conditions [Sambrook, J., et Molecular Cloning: A Laboratory Manual, Second Cold Spring Harbor Laboratory Press, Cold Edition. Double stranded DNA was Spring Harbor (1989)]. radiolabelled with 32P-Alpha-dCTP using a Pharmacia random priming labelling kit according to the manufacturer's To improve the labelling reaction cosmid instructions. DNA was digested with EcoRI prior to radiolabelling. test strips were hybridized without prehybridization in 10 plastic bags containing 6 x SSCP (saline, sodium citratephosphate buffer) and 0.01 % sodium dodecyl sulfate (SDS) Only target-specific binding to the for 16 hrs. polypropylene membranes was observed, eliminating the need for a prehybridization step. The specific activity 15 of the radiolabelled probes was adjusted to 5 \times 10 6 cpm/ml hybridization solution. After hybridization the test strips were washed in 2 x SSCP, 0.01 % SDS for 20 min. A variety of hybridization and wash temperatures was employed, as hereinafter described. Autoradiograms were 20 developed after 5 minutes to 6 hours exposure at -70° C. The resulting signals on the test strips were evaluated visually.

As expected, the synthetic (CAG), [SEQ ID NO:28] oligonucleotide probe hybridized specifically at 60°C to three rows of the array. These corresponded to the oligonucleotide repeats (CGT), [SEQ ID NO:57], (GTC), [SEQ ID NO:69], and (TCG), [SEQ ID NO:84], respectively.

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Using double-stranded DNA of 200 bp and 800 bp containing a (CTG)₁₁ [SEQ ID NO:29] or a (CAG)₁₀ [SEQ ID NO:30] repeat resulted in a pattern of 6 bands corresponding to (ACG)₇ [SEQ ID NO:37], (CGA)₇ [SEQ ID NO:54], (CGT)₇ [SEQ ID NO:57], (GAC)₇ [SEQ ID NO:63], (GTC)₇ [SEQ ID NO:75], and (TCG)₇ [SEQ ID NO:84] - i.e., the sense and antisense orientations. Differences in the signal intensity were observed between the various triplet-representing lanes.

Using the 3 kb PCR fragment containing a combined repeat (GCA)₈+(GCG)₄ [SEQ ID NO:31] to probe the test strips resulted in a complete set of six oligonucleotides - (ACG)₇ [SEQ ID NO:37], (CGA)₇ [SEQ ID NO:54], (CGT)₇ [SEQ ID NO:57], (GAC)₇ [SEQ ID NO:63], (GTC)₇ [SEQ ID NO:75], and (TCG)₇ [SEQ ID NO:84]. This set represents the six different frame shifts for the (GCA)₈ [SEQ ID NO:94] repeat. Additionally, the signals found with (CCG)₇ [SEQ ID NO:52], (CGC)₇ [SEQ ID NO:55], and (GCC)₇ [SEQ ID NO:67] were evident for the 3'--->5' directed frame of the (GCG)₄ [SEQ ID NO:95] repeat. No signal was detected under these conditions for the reversed direction indicated by (GGC)₇ [SEQ ID NO:71], (GCG)₇ [SEQ ID NO:68], and (CGC)₇ [SEQ ID NO:55].

Using cosmid MDY2 with an insert size of 31 kb as a probe, a band pattern was observed indicative of the and (CCT)_n (CAG), (GCC)_n, repeats, of presence For the (CCT)_n, only one direction of the respectively. oligonucleotide frame as represented by (CCT), [SEQ ID NO:53], (CTC), [SEQ ID NO:59], and (TCC), [SEQ ID NO: 83] was found to hybridize. A search of 11,613 nt available sequence information (GenBank accession L00727) of cosmid MDY2 revealed the presence of all types of triplet repeats identified by the test strip (Table 3). repeated triplet numbers vary from 3 for the CCT and GCG type repeats to 11 for the CTG repeat..

TABLE 3

Position
(Nucleotide #)

809-817

(CCT)₃

8,172-8,180

(CCT)₃

9,093-9,101

(GGA)₃

10,364-10,372

(GCC)₃

10,677-10,709

(SEQ ID NO:29]

The influence of temperature on the STR detection

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was evaluated by hybridizing cosmid 22.3 at 40°C, 50°C, and 60°C to the test strips. The strips were then washed used for the respective temperatures the At 40°C, a band pattern was obtained hybridizations. indicative of (CA)_n, (ACC)_n, (CCT), and (GCC), type repeats, respectively (Fig. 2A); for the triplet repeats, only one set of signals representing one direction of the oligonucleotide frame was observed. The pattern at 50°C was also specific for $(CA)_n$, $(ACC)_n$, and $(GCC)_n$ type repeats (Fig. 2B); however, unlike the pattern at 40°C the signals representing a $(TCC)_n$ type repeat disappeared and additional bands indicative of a (CGT), type repeat occurred; again, for the trinucleotide repeats only one signals representing one direction of set of respective oligonucleotide frame was found. At 60°C only representing (CA), and (CCG), type repeats signals Under hybridization these 2C). (Fig. persisted conditions a full set of the expected 6 bands evident for a (CCG), type repeat was observed.

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All types of STRs indicated by the test strips were found to be present in 34,977 bp sequence available from cosmid 22.3 (Fig. 3). They range from a single repeat unit of (CCT)₃ and (AGC)₃ up to 18 repeat units for a (TG)-dinucleotide repeat. There was no unspecific hybridization signal observed. AT-rich repeats also occurring in the sequence in three or less repeat units were not detected by the test strips under the hybridization conditions used.

The array was designed to represent trinucleotide repeats by all three possible frames in 3'-->5' direction, as well as in the reverse direction. Thus, using single stranded DNA a complementary sequence to a given trinucleotide repeat should result in three signals on the test strips; using double stranded DNA six respective bands for a given repeat should occur. For the four dinucleotide repeats only one frame was used for each type. Using this reverse blotting system, the

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obtained band pattern provided qualitatively the precise identification of previously known STRs in DNA samples of various complexities between 21 bp - 34,977 bp. Moreover, there was no random or cross hybridization to unspecific sequences observed. Based on the Tm and size of the STRs as well as possible influences by flanking sequences, varying the hybridization stringency can enhance the specificity.

art can readily ascertain the essential characteristics of the invention and, without departing from the spirit and scope thereof, can adapt the invention to various usages and conditions. Changes in form and substitution of equivalents are contemplated as circumstances may suggest or render expedient, and although specific terms have been employed herein, they are intended in a descriptive sense and not for purposes of limitation.

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SEQUENCE LISTING

```
(1) GENERAL INFORMATION:
 5
           (i) APPLICANT: Beckman Instruments, Inc.
10
          (ii) TITLE OF INVENTION: OLIGONUCLEOTIDE REPEAT ARRAYS
          (iii) NUMBER OF SEQUENCES: 95
15
           (iv) CORRESPONDENCE ADDRESS:
                 (A) ADDRESSEE: Robbins, Berliner & Carson
                 (B) STREET: 201 North Figueroa Street, Suite 500
                 (C) CITY: Los Angeles
                 (D) STATE: CA
20
                 (E) COUNTRY: USA
                 (F) ZIP: 90012
            (v) COMPUTER READABLE FORM:
                 (A) MEDIUM TYPE: Floppy disk
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                 (B) COMPUTER: IBM PC compatible
                 (C) OPERATING SYSTEM: PC-DOS/MS-DOS
                 (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
           (vi) CURRENT APPLICATION DATA:
30
                 (A) APPLICATION NUMBER:
                 (B) FILING DATE:
                 (C) CLASSIFICATION:
         (viii) ATTORNEY/AGENT INFORMATION:
35
                 (A) NAME: Spitals, John P.
                 (B) REGISTRATION NUMBER: 29,215
                 (C) REFERENCE/DOCKET NUMBER: 5727-110
           (ix) TELECOMMUNICATION INFORMATION:
                 (A) TELEPHONE: (213) 977-11001
40
                 (B) TELEFAX: (213) 977-1003
       (2) INFORMATION FOR SEQ ID NO:1:
 45
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 12 base pairs
                 (B) TYPE: nucleic acid
                 (C) STRANDEDNESS: single
 50
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: Other nucleic acid
.55
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
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                  (A) LENGTH: 12 base pairs
                  (B) TYPE: nucleic acid
                  (C) STRANDEDNESS: single
 65
                  (D) TOPOLOGY: linear
            (ii) MOLECULE TYPE: Other nucleic acid
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15	(ii) MOLECULE TYPE: Other nucleic acid	
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30	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ 1D NO:4:	
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45	(ii) MOLECULE TYPE: Other nucleic acid	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	15
·55	(2) INFORMATION FOR SEQ ID NO:6: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
60	(ii) MOLECULE TYPE: Other nucleic acid	
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

-28-

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25	(ii) MOLECULE TYPE: Other nucleic acid	
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	(ii) MOLECULE TYPE: Other nucleic acid	
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	וווווווווו ווווווווווווווווווווווווווו	15
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	(ii) MOLECULE TYPE: Other nucleic acid	
60		
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C F	ааааааааа аааааааааа А	21
65	(2) INFORMATION FOR SEQ ID NO:11:	
70	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

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	(ii) MOLECULE TYPE: Other nucleic acid	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
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15	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
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^-	GGGGGGGGG GGGGGGGG G	٤١
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30	(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
35	(11) HOLLOGE THE Street Hastons do to	
55	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
	THITTITI THITTITI I	21
40	(2) INFORMATION FOR SEQ ID NO:14:	۵,
45	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
55	AACAACAACA ACAACAACAA C	21
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60	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
65	(ii) MOLECULE TYPE: Other nucleic acid	
70	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
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(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
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(ii) MOLECULE TYPE: Other nucleic acid	
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TGCTGCTGCT GCTGCTGC	18
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(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
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TGCTGCTGCT GCTGTGTGT TGTGTTATTA TTATTA	36
(2) INFORMATION FOR SEQ ID NO:19: (i) SEQUENCE CHARACTERISTICS: (A) 1 FNGTH: 24 base pairs	
(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:	
TGGTGGTGGT GGTGATGATGA	24
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(ii) MOLECULE TYPE: Other nucleic acid

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•	TCGTCGTCGT CGTCGTCG	18
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	(ii) MOLECULE TYPE: Other nucleic acid	
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	GGCGGCGGCG GCGGC	15
25	(2) INFORMATION FOR SEQ ID NO:22:	
° 30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
35		
	(xi) SEQUENCE DESCRIPTION: SEQ 1D NO:22:	
40	CGCGCGCGGA CGACGACGAC	20
- 0	(2) INFORMATION FOR SEQ ID NO:23:	
45	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
55	TCATCATCAT CATAATAATA ATAATACTAC TACTAC	36
	(2) INFORMATION FOR SEQ ID NO:24:	
60	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
65	(ii) MOLECULE TYPE: Other nucleic acid	
70	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
, 0	CCACCACCAC CACAACAACA ACAA	24

	(2) INFORMATION FOR SEQ TO NO.23:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
15	דוכווכווכו וכוכוכוכו וכוכוככוככ וככוככ	36
	(2) INFORMATION FOR SEQ 1D NO:26:	•
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: Other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
J 0	GAAGAAGAAG AAGAGAGAGA GAGAGGA GGAGGA	36
	(2) INFORMATION FOR SEQ ID NO:27:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: Other nucleic acid	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	
	GCCGCCGCCG CCGCAGCAGC AGCA	24
50	(2) INFORMATION FOR SEQ ID NO:28: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs	
·55	(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
60	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
	CAGCAGCAGC AGCAGCAGCA G	21
65	(2) INFORMATION FOR SEQ ID NO:29:	
70	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 33 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
	(D) TOPOLOGY: Linear	

•	i	i١	MOI	FCILLE	TYPE -	Other	nucleic	acid
	•		1100	LLULE	I I FE.	ULHEI	HULLEIL	acıu

5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
	CTGCTGCTGC TGCTGCTGCT GCTGCTGCTG	33
10	(2) INFORMATION FOR SEQ ID NO:30:	
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	•
20		
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
	CAGCAGCAGC AGCAGCAGCAG	30
25	(2) INFORMATION FOR SEQ ID NO:31:	
30	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: Other nucleic acid	•
35		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	
40	GCAGCAGCAG CAGCAGCAGC AGCAGCGGCG GCGGCG	36
	(2) INFORMATION FOR SEQ ID NO:32:	·
45	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
55	ACACACACA ACACACACAC	20
	(2) INFORMATION FOR SEQ ID NO:33:	
60	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
65	(ii) MOLECULE TYPE: Other nucleic acid	
70	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
, ,	AAGAAGAAGA AGAAGAAGAA G	21

	(2) INFORMATION FOR SEQ ID NO:34:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:	
15	AATAATAATA ATAATAATAA T	21
	(2) INFORMATION FOR SEQ 1D NO:35:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: Other nuclei: acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:	
	ACAACAACAA CAACAACAAC A	21
	(2) INFORMATION FOR SEQ ID NO:36:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: Other nucleic acid	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:	
	ACCACCACCA CCACCACCAC C	
50	(2) INFORMATION FOR SEQ ID NO:37: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid	
55	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
60	(xi) SEQUENCE DESCRIPTION: SEQ 10 NO:37:	
	ACGACGACGA CGACGACGAC G	21
65	(2) INFORMATION FOR SEQ ID NO:38:	
5 0	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid	
70	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	

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	(ii) MOLECULE TYPE: Other nucleic acid	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:	
	ACTACTACTA CTACTACTAC T	21
	(2) INFORMATION FOR SEQ ID NO:39:	
10 15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
13	(ii) MOLECULE TYPE: Other nucleic acid	
	(11) Modecute 1172: Other Indicate acid	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:	
	AGAAGAAGAA GAAGAAGAAG A	21
25	(2) INFORMATION FOR SEQ ID NO:40:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
35		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:	
40	AGCAGCAGCA GCAGCAGCAG C	
40	(2) INFORMATION FOR SEQ ID NO:41:	
45	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(x1) SEQUENCE DESCRIPTION: SEQ ID NO:41:	
55	AGGAGGAGGA GGAGGAGGAG G	21
	(2) INFORMATION FOR SEQ ID NO:42:	
60	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
65	(ii) MOLECULE TYPE: Other nucleic acid	
70	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:	
	AGTAGTAGTA GTAGTAGTAG T	21

	(2) INFORMATION FOR SEQ ID NO:45:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:	
15	ATAATAATAATAATAA	21
	(2) INFORMATION FOR SEQ ID NO:44:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: Other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:	
30	ATCATCATCA TCATCATCAT C	
	(2) INFORMATION FOR SEQ ID NO:45:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: Other nucleic acid	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:	
	ATGATGATGA TGATGAT G	21
50	(2) INFORMATION FOR SEQ ID NO:46:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
.55	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
60	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:	
	ATTATTATTA TTATTATTAT T	21
65	(2) INFORMATION FOR SEQ ID NO:47:	
Ų J	(i) SEQUENCE CHARACTERISTICS:	
70	(A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

70

	(ii) MOLECULE TYPE: Other nucleic acid	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:	
	CAACAACAAC AACAACAACA A	21
	(2) INFORMATION FOR SEQ ID NO:48:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
15	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:	
	CACCACCACC ACCACCACCA C	
25	(2) INFORMATION FOR SEQ ID NO:49:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: rucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
35		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:	
	CATCATCATC ATCATCATCA T	21
40	(2) INFORMATION FOR SEQ ID NO:50:	
45	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:	
.55	CCACCACCAC CACCACCAC A	
	(2) INFORMATION FOR SEQ ID NO:51:	
60	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
65	(ii) MOLECULE TYPE: Other nucleic acid	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:
CGCGCGCGCG CGCGCGCGCG 20

	(2) INFORMATION FOR SEQ ID NO:52:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:	
15	CCGCCGCCGC CGCCGCCGC G	21
	(2) INFORMATION FOR SEQ ID NO:53:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: Other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:	
	CCTCCTCCTC CTCCTCCC T	21
	(2) INFORMATION FOR SEQ ID NO:54:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: Other nucleic acid	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:	
	CGACGACGAC GACGACG A	
50	(2) INFORMATION FOR SEQ ID NO:55: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid	
55	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
0.	(ii) MOLECULE TYPE: Other nucleic acid	
60	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:	
	CGCCGCCGCC GCCGCCGCC C	21
65	(2) INFORMATION FOR SEQ ID NO:56:	
70	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
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	(11) Hotelste 111 at a sile in the sile in	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:	
	CGCCGCCGC GCCGCCGCC G	21
	(2) INFORMATION FOR SEQ 1D NO:57:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
15	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:	
	CGTCGTCGTC GTCGTCGTCG T	21
25	(2) INFORMATION FOR SEQ ID NO:58:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
35		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:	
40	CTACTACTAC TACTACTACT A	
40	(2) INFORMATION FOR SEQ ID NO:59:	
45	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:	
.55	стестестее тестестеет с	21
	(2) INFORMATION FOR SEQ ID NO:60:	
60	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
65	(ii) MOLECULE TYPE: Other nucleic acid	
70	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:	
, 0	CTGCTGCTGC TGCTGCT G	21

	(2) INFORMATION FOR SEQ ID NO:61:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:61:	
15	כדוכדוכדוכ דוכדוכדוכד ד	21
	(2) INFORMATION FOR SEQ ID NO:62:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: Other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:	
	GAAGAAGAAG AAGAAGAAGA A	
	(2) INFORMATION FOR SEQ ID NO:63:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: Other nucleic acid	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:	
	GACGACGACG ACGACGACGA C	21
50	(2) INFORMATION FOR SEQ ID NO:64: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid	
-55	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
60	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:	
	GAGGAGGAGG AGGAGGAGGA G	21
65	(2) INFORMATION FOR SEQ ID NO:65:	
70	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

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(ii) MOLECULE TYPE: Other nucleic acid
 5
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:
                                                                               21
      GATGATGATG ATGATGATGA T
      (2) INFORMATION FOR SEQ ID NO:66:
10
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 21 base pairs
                 (B) TYPE: nucleic acid
                 (C) STRANDEDNESS: single
15
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: Other nucleic acid
20
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:
      GCAGCAGCAG CAGCAGCAGC A
25
       (2) INFORMATION FOR SEQ ID NO:67:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 21 base pairs
                 (B) TYPE: nucleic acid
30
                 (C) STRANDEDNESS: single
                 (D) TOPOLOGY: Linear
           (ii) MOLECULE TYPE: Other nucleic acid
35
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:
                                                                               21
       GCCGCCGCCG CCGCCGCCCC C
40
       (2) INFORMATION FOR SEQ ID NO:68:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 21 base pairs
                 (B) TYPE: nucleic acid
(C) STRANDEDNESS: single
45
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: Other nucleic acid
50
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:
.55
       GCGGCGGCGG CGGCGGCGGC G
       (2) INFORMATION FOR SEQ ID NO:69:
            (1) SEQUENCE CHARACTERISTICS:
                  (A) LENGTH: 21 base pairs
 60
                  (B) TYPE: nucleic acid
                  (C) STRANDEDNESS: single
                  (D) TOPOLOGY: linear
 65
            (ii) MOLECULE TYPE: Other nucleic acid
            (xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:
 70
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GCTGCTGCTG CTGCTGCTGC T

	(2) INFORMATION FOR SEQ ID NO:70:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:70:	
15	GGAGGAGGAG GAGGAGGAG A	21
	(2) INFORMATION FOR SEQ ID NO:71:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: Other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:	
	GGCGGCGCGC GCGCGCGC C	
	(2) INFORMATION FOR SEQ 1D NO:72:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: Other nucleic acid	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:72:	
	AGAGAGAGA AGAGAGAGAG	20
50	(2) INFORMATION FOR SEQ ID NO:73: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs	
·55	(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
60	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:73:	
	GGTGGTGGTG GTGGTGGTGG T	21
65	(2) INFORMATION FOR SEQ ID NO:74:	
70	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEONESS: single (D) TOPOLOGY: linear	

21

(ii) MOLECULE TYPE: Other nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO:74: 5 21 GTAGTAGTAG TAGTAGTAGT A (2) INFORMATION FOR SEQ ID NO:75: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 15 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:75: GTCGTCGTCG TCGTCGTCGT C (2) INFORMATION FOR SEQ ID NO:76: 25 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid 30 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid 35 (xi) SEQUENCE DESCRIPTION: SEQ ID 40:76: 21 GTGGTGGTGG TGGTGGTGGT G 40 (2) INFORMATION FOR SEQ ID NO:77: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs 45 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid 50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:77: 21 ·55 GTTGTTGTTG TTGTTGTTGT T (2) INFORMATION FOR SEQ ID NO:78: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs 60 (8) TYPE: nucleic acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(C) STRANDEDNESS: single

70 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:78:

(2) INFORMATION FOR SEW ID NO:19:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:79:	
TACTACTACT ACTACTACTA C	
(2) INFORMATION FOR SEQ ID NO:80:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:80:	
TAGTAGTAGT AGTAGTAGTA G	21
(2) INFORMATION FOR SEQ ID NO:81:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID.NO:81:	
TATTATTATT ATTATTATTA T	21
(2) INFORMATION FOR SEQ ID NO:82: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs	
(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:82:	
TCATCATCAT CATCATCATC A	21
(2) INFORMATION FOR SEQ ID NO:83:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO:79: TACTACTACT ACTACTACTA C (2) INFORMATION FOR SEQ ID NO:80: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO:80: TAGTAGTAGT AGTAGTAGTA G (2) INFORMATION FOR SEQ ID NO:81: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO:81: TATTATTATT ATTATTATTA T (2) INFORMATION FOR SEQ ID NO:82: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid (xi) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO:82: TCATCATCAT CATCATCATC A (2) INFORMATION FOR SEQ ID NO:83: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid

	(11) MOLECULE 11FE. Other Mucters acid	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:83:	
	TECTCCTCCT CCTCCTCC C	
10	(2) INFORMATION FOR SEQ ID NO:84:	
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
20		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:84:	
25	TCGTCGTCGT CGTCGTCGTC G	2
	(2) INFORMATION FOR SEQ ID NO:85:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
35	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:85:	
40	теттеттетт сттеттетте т	2
	(2) INFORMATION FOR SEQ ID NO:86:	
45	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:86:	
55	TGATGATGAT GATGATGATG A	2
	(2) INFORMATION FOR SEQ ID NO:87:	
60	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
65	(ii) MOLECULE TYPE: Other nucleic acid	
70	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:87:	

TGCTGCTGCT GCTGCTGCTG C

(2) INFORMATION FOR SEQ ID NO:88:

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5 -	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:88:	
15	TEGTEGTEGT GETEGTEGTE G	21
	(2) INFORMATION FOR SEQ ID NO:89:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: Other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:89:	
	(2) INFORMATION FOR SEQ ID NO:90:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
40	(D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid	
45	(x1) SEQUENCE DESCRIPTION: SEQ ID NO:90:	
	TTATTATTAT TATTATTATT A	
50	(2) INFORMATION FOR SEQ ID NO:91: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs	
55	(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
60		•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:91:	
	דוכוזכווכן וכווכווכון כ	21
65	(2) INFORMATION FOR SEQ ID NO:92:	
70	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

(ii) MOL	ECULE	TYPE:	Other	nucl	leic	acid
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5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:92:	
	TIGITGITGT TGTIGITGIT G	
10	(2) INFORMATION FOR SEQ ID NO:93:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
15	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
20		
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:93:	
	כזכזכזכזכז כזכזכזכזכז	20
25	(2) INFORMATION FOR SEQ ID NO:94:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
30	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
35		
	(x1) SEQUENCE DESCRIPTION: SEQ ID NO:94:	•
40	GCAGCAGCAG CAGCAGCAGC AGCA	24
	(2) INFORMATION FOR SEQ ID NO:95:	
45	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 12 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:95:	
	GCGGCGGCGG CG	12

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WHAT IS CLAIMED IS:

1. Apparatus for identifying tandem nucleotide repeats in a sample containing same, comprising:

a solid support; and

oligonucleotides fixedly plurality of attached thereto to form the an array, oligonucleotides defining a set of tandem nucleotide repeats including sequences complementary to tandem repeats in the sample, the nucleotide establishing a pattern such that identity of a tandem nucleotide repeat in the sample may be ascertained by location in the pattern of the nucleotide complementary tandem repeat upon hybridization of the sample to the array.

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- 2. Apparatus according to claim 1, wherein the array comprises a linear sequence of oligonucleotides.
- 3. Apparatus according to claim 1, wherein the 20 array comprises a two-dimensional pattern.
- 4. Apparatus according to claim 1, wherein the oligonucleotides have the formula (Nmer), in which N is an integer greater than 1 and represents the number of nucleotides in a repeat pattern and n is an integer from two to about 50.
- 5. Apparatus according to claim 4, wherein the set of tandem nucleotide repeats comprises a complete set of 30 (3mer), repeats excluding homopolymers.
 - 6. Apparatus according to claim 1, wherein the set of tandem nucleotide repeats comprises a complete set of (2mer), repeats excluding homopolymers.

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7. Apparatus according to claim 1, wherein the set of tandem nucleotide repeats comprises a set of oligonucleotides of the formula

 $[(Nmer)_n(Mmer)_m)]_x$

5 or

[(Nmer)_n(Mmer)_m(Pmer)_p]_x

in which each of N, M and P is independently selected from the set of integers greater than 1 and represent the number of nucleotides in a repeat pattern;

each of n, m and p is independently selected from the set of integers; and

x is an integer, with the proviso that x(Nn + Mm) or x(Nn + Mm + Pp) is between 4 and about 100.

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- 8. Apparatus according to claim 1, wherein the solid support comprises a material selected from glass and plastic films.
- 9. Apparatus according to claim 8, wherein the solid support comprises polypropylene.
 - 10. A method for identifying tandem nucleotide repeats in a sample containing same, comprising:

bringing the sample into contact under hybridization conditions with an array comprising a plurality of oligonucleotides fixedly attached to a solid support, the oligonucleotides defining a set of tandem nucleotide repeats including sequences complementary to tandem nucleotide repeats in the sample, the array establishing a predetermined pattern; and

identifying the tandem nucleotide repeats in the sample by determining location in the pattern of the complementary tandem nucleotide repeats to which the tandem nucleotide repeats are hybridized. WO 95/30774 PCT/US95/04899

- 11. A method according to claim 10, wherein the sample is labelled prior to bringing it into contact with the array.
- 12. A method according to claim 11, wherein the label is selected from the group consisting of radioactive labels and fluorescent labels.
- 13. A method according to claim 10, wherein the oligonucleotides have the formula (Nmer), in which N is an integer greater than 1 and represents the number of nucleotides in a repeat pattern and n is an integer from two to about 50.
- 14. A method according to claim 10, wherein the set of tandem nucleotide repeats comprises a complete set of (3mer), repeats excluding homopolymers.
- 15. A method according to claim 10, wherein the set 20 of tandem nucleotide repeats comprises a complete set of (2mer)_n repeats excluding homopolymers.
- 16. A method according to claim 10, wherein the set of tandem nucleotide repeats comprises a set of 25 oligonucleotides of the formula

 $[(Nmer)_n(Mmer)_m)]_x$

or

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[(Nmer)_n(Mmer)_m(Pmer)_p]_x

in which each of N, M and P is independently selected from the set of integers greater than 1 and represent the number of nucleotides in a repeat pattern;

each of n, m and p is independently selected from the set of integers; and

35 x is an integer, with the proviso that x(Nn + Mm) or x(Nn + Mm + Pp) is between 4 and about 100.

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17. A method for determining size of tandem nucleotide repeats in a sample containing same, comprising:

bringing the sample into contact under hybridization conditions with an array comprising a plurality of oligonucleotides fixedly attached to a solid support, the oligonucleotides defining a set of tandem nucleotide repeats including sequences complementary to tandem nucleotide repeats in the sample, the array establishing a predetermined pattern;

identifying the tandem nucleotide repeats in the sample by determining location in the pattern of the complementary tandem nucleotide repeats to which the tandem nucleotide repeats are hybridized to form hybrids;

selectively dissociating the hybrids to elute the tandem nucleotide repeat from the support; and sizing the eluted tandem nucleotide repeats.

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18. A method according to claim 17, wherein the eluted tandem nucleotide repeats are sized by gel electrophoresis against DNA size markers.

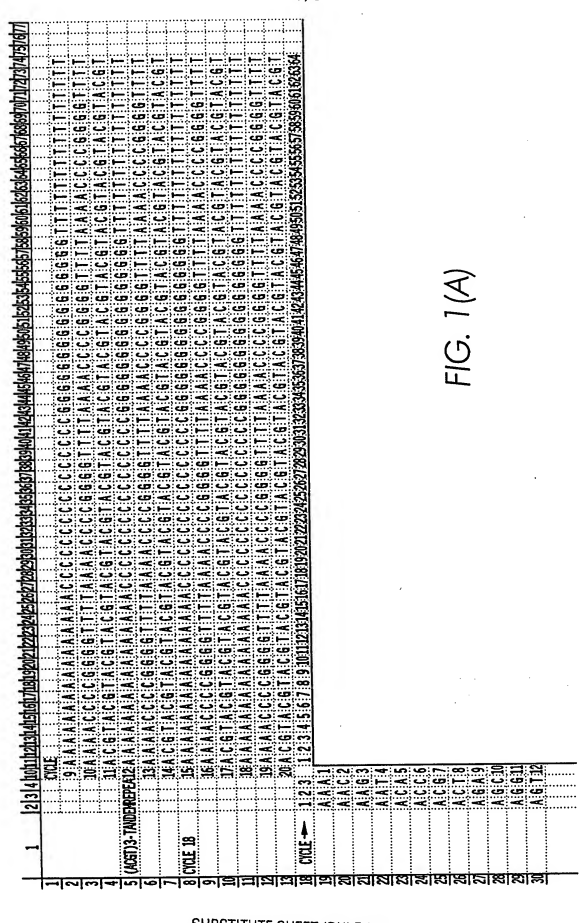
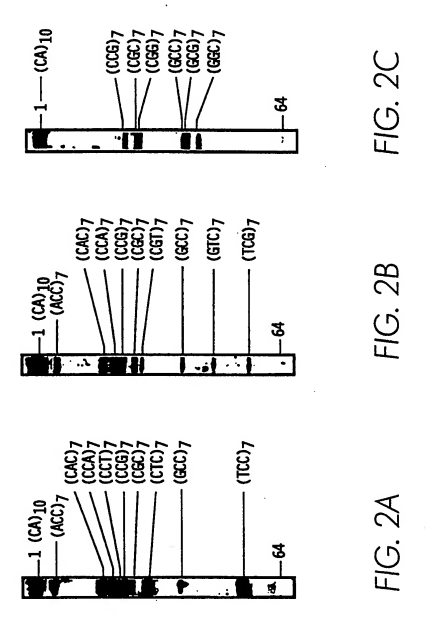


FIG. 1(B)

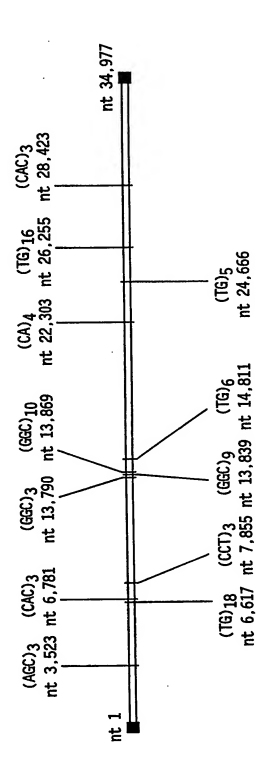


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INTERNATIONAL SEARCH REPORT

Inten all Application No PCT/US 95/04899

A. CLASS IPC 6	SIFICATION OF SUBJECT MATTER C12Q1/68			
According t	to International Patent Classification (IPC) or to both national classi	ification and IPC		
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Minimum d	documentation searched (classification system followed by classificat C120	tion symbols)	· \	
IFC 0	C12Q			
Documental	tion searched other than minimum documentation to the extent that	such documents are included in the fields so	earched	
	·			
Electronic data base consulted during the international search (name of data base and, where practical, search terms used)				
C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category *	Citation of document, with indication, where appropriate, of the re	elevant passages	Relevant to claim No.	
Υ	WO,A,92 10588 (AFFYMAX TECH IIV) 2 1992 see the whole document	25 June	1-18	
Y	WO,A,89 10977 (ISIS INNOVATION) 1 November 1989 cited in the application see the whole document	16	1-18	
Υ	WO,A,93 17126 (PUBLIC HEALTH RESS OF) 2 September 1993 see the whole document	EARCH INST	1-18	
		-/		
Further documents are listed in the continuation of box C. Patent family members are listed in annex.				
"A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but		"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family		
Date of the actual completion of the international search 4 September 1995		Date of mailing of the international search report 1 9. 09. 95		
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016		Authorized officer Molina Galan, E		

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INTERNATIONAL SEARCH REPORT

Inter: 1al Application No PCT/US 95/04899

Category *	tion) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Caugury	Company of monumental community control of the community control of the control o	
Y	HUMAN MOLECULAR GENETICS, vol. 3, no. 4, April 1994 OXFORD GB, pages 599-605, ARMOUR ET AL. 'Isolation of human simple repeat loci by hybridisation selection' see the whole document	1-18
Y	NUCLEIC ACIDS RESEARCH, vol. 21, no. 16, 1993 OXFORD GB, pages 3911-3912, KARAAGYOZOV ET AL. 'Construction of random small-insert genomic libraries highly enriched for simple sequence repeats' see the whole document	1-18
A	GENETIC ANALYSIS TECHNIQUES AND APPLICATIONS, vol. 10, no. 1, 1993 NEW YORK US, pages 2-5, IIZUKA ET AL. 'Selective isolation of highly polymorphic (C-A)n * (D-G)n microsatellites by stringent hybridisation' cited in the application	
P,X	NUCLEIC ACIDS RESEARCH, vol. 22, no. 9, 11 May 1994 OXFORD GB, pages 1701-1704, WEHNERT ET AL. 'A rapid scanning strip for tri- and dinucleotide short tandem repeats' see the whole document	1-18